

# UC Davis

## UC Davis Previously Published Works

### Title

Vectors with autonomy: what distinguishes animal-mediated nutrient transport from abiotic vectors?

### Permalink

<https://escholarship.org/uc/item/3x63w60w>

### Journal

Biological reviews of the Cambridge Philosophical Society, 94(5)

### ISSN

1464-7931

### Authors

McInturf, Alexandra G  
Pollack, Lea  
Yang, Louie H  
et al.

### Publication Date

2019-10-01

### DOI

10.1111/brv.12525

Peer reviewed

**1 Vectors with autonomy: what distinguishes animal-**  
**2 mediated nutrient transport from abiotic vectors?**

3

4 Alexandra G. McInturf<sup>1,\*†</sup>, Lea Pollack<sup>2,†</sup>, Louie H. Yang<sup>3</sup> and Orr  
5 Spiegel<sup>4</sup>

6

7 *<sup>1</sup>Department of Wildlife, Fish, and Conservation Biology, University of*  
8 *California, One Shields Avenue, Davis, CA, 95616, USA*

9 *<sup>2</sup> Department of Environmental Science and Policy, University of California,*  
10 *One Shields Avenue, Davis, CA, 95616, USA*

11 *<sup>3</sup>Department of Entomology and Nematology, University of California, Davis,*  
12 *One Shields Avenue, Davis, CA, 95616, USA*

13 *<sup>4</sup>Sherman Building, School of Zoology, Faculty of Life Sciences, Tel Aviv*  
14 *University, Tel Aviv, 69978, Israel*

15

16 \*Author for correspondence (E-mail: amcinturf@ucdavis.edu; Tel.: 513-504-  
17 8939).

18 <sup>†</sup>Equal first authors.

19

20 ABSTRACT

21       Animal movements are important drivers of nutrient redistribution that  
22 can affect primary productivity and biodiversity across various spatial scales.

Recent work indicates that incorporating these movements into ecosystem models can enhance our ability to predict the spatio-temporal distribution of nutrients. However, the role of animal behaviour in animal-mediated nutrient transport (i.e. active subsidies) remains under-explored. Here we review the current literature on active subsidies to show how the behaviour of active subsidy agents makes them both ecologically important and qualitatively distinct from abiotic processes (i.e. passive subsidies). We first propose that animal movement patterns can create similar ecological effects (i.e. press and pulse disturbances) in recipient ecosystems, which can be equal in magnitude to or greater than those of passive subsidies. We then highlight three key behavioural features distinguishing active subsidies. First, organisms can transport nutrients counter-directionally to abiotic forces and potential energy gradients (e.g. upstream). Second, unlike passive subsidies, organisms respond to the patterns of nutrients that they generate. Third, animal agents interact with each other. The latter two features can form positive- or negative-feedback loops, creating patterns in space or time that can reinforce nutrient hotspots in places of mass aggregations and/or create lasting impacts within ecosystems. Because human-driven changes can affect both the space-use of active subsidy species and their composition at both population (i.e. individual variation) and community levels (i.e. species interactions), predicting patterns in nutrient flows under future modified environmental conditions depends on understanding the behavioural mechanisms that underlie active subsidies and variation among agents'

46contributions. We conclude by advocating for the integration of animal  
47behaviour, animal movement data, and individual variation into future  
48conservation efforts in order to provide more accurate and realistic  
49assessments of changing ecosystem function.

50

51*Key words:* animal behaviour, behavioural ecology, energy flow, mobile links,  
52movement ecology, nutrient cycling, spatial subsidies.

53

## 54CONTENTS

55I. Introduction: animals as important vectors of nutrient transport

56II. The quantitative importance of active subsidies

57III. Features that distinguish active from passive subsidies

58     (1) Counter-gradient and cross-habitat transport

59     (2) Agents respond to the nutrient distribution patterns that they  
60     generate

61     (3) Interactions among agents affect deposition patterns

62IV. Individual differences as a source of variation

63V. Conservation implications

64VI. Conclusions

65VII. Acknowledgements

66VIII. References

67

## 68I. INTRODUCTION: ANIMALS AS IMPORTANT VECTORS OF NUTRIENT 69TRANSPORT

70       The concentration of nutrients across multiple spatio-temporal scales  
71establishes the foundation of ecosystem productivity and subsequent  
72diversity within and across habitats. Geological processes such as tectonic  
73movement and sedimentation determine the underlying distribution of  
74essential resources (e.g. nitrogen, phosphorous, and carbon) that influence  
75primary productivity and biomass (Cook & McElhinny, 1979; Vitousek, 2004;  
76Elser *et al.*, 2007; LeBauer & Treseder, 2008; Vitousek *et al.*, 2013). This  
77initial distribution is modified by environmental and physical factors ('passive  
78subsidies'; Earl & Zollner, 2014), including wind, current, gravity and erosion  
79(Zhao & Running, 2010; Cleveland *et al.*, 2013; Houlton & Morford, 2015;  
80Morford, Houlton & Dahlgren, 2016). In addition to these abiotic processes,  
81biotic vectors further redistribute nutrients through various mechanisms,  
82mostly *via* animal movement (Fig. 1). Specifically, animals can displace  
83resources, or serve as 'mobile links' (Jeltsch *et al.*, 2013), within and among  
84ecosystems, generating nutrient inputs that are referred to as 'active  
85subsidies' (Earl & Zollner, 2014). Active subsidies often differ in their physical  
86form (nutrient composition, lability, etc.), which can influence the pathways  
87by which these nutrient influxes enter ecosystems (Marcarelli *et al.*, 2011).  
88Furthermore, animal vectors exhibit diverse behaviours that influence  
89movement patterns and how nutrients are distributed. Ultimately, these  
90behaviours allow active subsidies to be quantitatively important in

91ecosystem dynamics, but qualitatively different from passive subsidies in  
92their nutrient deposition patterns.

93       There is an extensive body of research demonstrating that animals  
94across taxa transport nutrients within and among ecosystems (e.g. Polis,  
95Anderson & Holt, 1997; Helfield & Naiman, 2001; Bauer & Hoyer, 2014;  
96Adame *et al.*, 2015). This literature shows that animals can redistribute large  
97masses of nutrients in the environment by (1) consuming and transporting  
98biomass (e.g. the deposition of waste products by migrating grazers such as  
99wildebeest (*Connochaetes taurinus*; Holdo *et al.*, 2007) and (2) serving as the  
100supply of biomass themselves (e.g. *via* predator consumption or  
101decomposition of wildebeest carcasses; Subalusky *et al.*, 2017). Yet, while  
102the impact of active subsidies in nutrient transportation has been  
103convincingly demonstrated in many studies, empirical work has historically  
104focused on the patterns of nutrient accumulation created by organism  
105movement, neglecting the processes by which animal vectors (directly or  
106indirectly) shape the ecosystem.

107       To address this limitation, recent work has explored how animal  
108movement decisions influence the distribution of resources in space and  
109time (e.g. Bauer & Hoyer, 2014; Earl & Zollner, 2017). This requires  
110consideration of the factors that drive an organism's behaviour, such as  
111characteristics of the external environment (e.g. initial nutrient distribution  
112or presence of other agents). A recent framework argues that exogenous  
113(external) factors such as mortality and competition risk, thermal

114conservation, and prey activity affect an animal's behaviour at multiple  
115hierarchical levels, from instantaneous decision-making to annual activity  
116patterns such as migration (McCann, Zollner & Gilbert, 2017). Collectively,  
117such factors interact with other drivers and constraints (such as the  
118organism's internal state, cognitive navigational capacity, and biomechanical  
119motion capacity) to shape each movement path (Nathan *et al.*, 2008). For  
120instance, wildebeest respond to external factors such as drying vegetation  
121through mass migration, and their collective movement ultimately results in  
122a relatively large nutrient influx into local river systems (Subalusky *et al.*,  
1232017). Predictions regarding future influx into rivers depend on the size of  
124the expected wildebeest population, the future environmental conditions  
125affecting their migration, and their consequent movement decisions. Thus, a  
126more mechanistic understanding of active subsidy transport is critical for  
127projecting alterations in nutrient patterns on the landscape, especially in  
128light of ongoing environmental change, which can alter community  
129composition (Barnosky *et al.*, 2012; Dirzo *et al.*, 2014), reduce population  
130sizes (Both *et al.*, 2006; Worm *et al.*, 2006) and affect the movements of  
131various species (Tucker *et al.*, 2018). Consequently, an increasing body of  
132literature now advocates modelling methods that consider animal movement  
133when predicting the ecological impact of active subsidies (e.g. individual-  
134based models, state-space models, hidden Markov models) (Earl & Zollner,  
1352017; Subalusky & Post, 2018).

136        Here, we review the quantitative importance of active subsidies and  
137their ability to generate substantial ecological effects (i.e. press and pulse  
138disturbances). We then offer a complementary behavioural perspective that  
139identifies three fundamental features that separate active subsidies from  
140passive subsidies. (1) First, active subsidies can move counter-directionally  
141to environmental gradients, transporting nutrients in directions that oppose  
142major abiotic forces like gravity, prominent wind flows, and prevailing  
143currents (e.g. salmon swimming upstream). (2) Second, active subsidy  
144agents can respond to the patterns of nutrients in their environment,  
145creating positive or negative feedbacks with the distribution of abiotic  
146resources. (3) Third, active subsidy agents respond to the presence and  
147behaviour of other organisms, creating positive or negative feedbacks with  
148aspects of the biotic environment, including other active subsidy agents.  
149Finally, we call attention to an area for future study by discussing the  
150potential impact of individual behavioural variation on active subsidy  
151distribution. With the support of several examples from the existing literature  
152that highlight these concepts, we conclude that understanding the  
153behavioural context of animal movement is essential for predicting and  
154conserving resource patterns formed by active subsidies in rapidly changing  
155environments.

156

## 157II. THE QUANTITATIVE IMPORTANCE OF ACTIVE SUBSIDIES



158       Recent work argues that the nutrient contributions from active  
159subsidies can be of similar magnitude to those of passive subsidies, and are  
160essential for many ecological systems (Earl & Zollner, 2017; Subalusky &  
161Post, 2018). Even small organisms can show behaviours (e.g. emergence  
162along the aquatic-terrestrial interface) that result in nutrient movements  
163that are substantially greater than those generated by abiotic forces (Yang &  
164Gratton, 2014). Similar evidence has been found in other systems with  
165varying vector species (see Fig. 2). These impacts are highly dependent on  
166the spatial and temporal scales of the subsidy's movement behaviour and its  
167interaction with prevailing passive nutrient-movement processes. For  
168example, populations of flying insects selectively follow prevailing winds,  
169effectively redistributing large amounts of biomass on a regional scale in  
170combination with abiotic processes (Hu *et al.*, 2017).

171       In addition to sheer magnitude, active subsidy inputs can act like  
172critical press or pulse perturbations within an ecosystem (defined below),  
173depending largely on the temporal heterogeneity of animal movements  
174(Bender, Case & Gilpin, 1984; Bauer & Hoyer, 2014; Allgeier, Burkepile &  
175Layman, 2017). Press disturbances are created by continuous movements  
176that lead to a sustained nutrient influx; these can occur when active subsidy  
177nutrient transport is relatively consistent in time, driven by repeated  
178patterns of activity (Wagner, Jones & Gordon, 2004; Fagan, Lutscher &  
179Schneider, 2007) (Fig. 3A). For example, Brazilian cave-dwelling bats  
180(*Hipposideros* and *Eonycteris* spp.) sustain entire cave ecosystems through

181their nightly roosting behaviour, since their guano provides nutrients to an  
182otherwise energy-poor system (Ferreira & Martins, 1999; Poulson & Lavoie,  
1832000; Fenolio *et al.*, 2006; Bird *et al.*, 2007; Kunz *et al.*, 2011). Parallel  
184examples also exist in marine environments (Williams *et al.*, 2018). By  
185contrast, pulse perturbations can occur when a less-common behaviour  
186results in the instantaneous alteration of active subsidy behaviour or  
187population density, and thus creates a flux of nutrients of large magnitude  
188and short duration (Fig. 3B, C). For example, the mass migration of  
189anadromous fishes can generate profound ecosystem impacts as they move  
190nutrients from the marine environment to aquatic and terrestrial systems  
191(Helfield & Naiman, 2001; Gende *et al.*, 2002; Varpe, Fiksen & Slotte, 2005;  
192Childress & McIntyre, 2015). Similar effects can result from the population  
193dynamics and movements of many other animals, including insects (Yang,  
1942004; Yang & Gratton, 2014; Hu *et al.*, 2017) and mammals (Roman &  
195McCarthy, 2010; Subalusky *et al.*, 2017). Depending on their frequency,  
196pulses can be cyclic (within a regular interval) or irregular. Although the  
197former is likely more predictable from the perspective of a consumer,  
198irregular pulses (such as locust outbreaks) can produce greater shifts from  
199baseline nutrient levels than mere seasonal changes (Fig. 3C). Thus,  
200although it can be easy to ignore nutrient fluxes driven by animal behaviour  
201as inconsistent or inconsequential when modelling community-level  
202dynamics, it is important to note that presses and pulses generated by active

subsidy input can be equally, if not more, instrumental to ecological processes as those of passive subsidies.

205

### 206 **III. FEATURES THAT DISTINGUISH ACTIVE FROM PASSIVE SUBSIDIES**

#### 207 **(1) Counter-gradient and cross-habitat transport**

208       Passive subsidies (e.g. erosion, water flow) often follow a gradient of  
209 potential energy, such as downstream water flow. While there are  
210 exceptional examples where passively transported nutrients move against a  
211 gradient of potential energy (e.g. wildfire, volcanic eruptions and upwelling),  
212 these are generally localized events. Organisms, on the other hand, can  
213 widely and selectively redistribute resources by moving them in directions  
214 and/or to distances unachievable by most passive agents ('counter-gradient  
215 transport'; Table 1). For instance, animals frequently exhibit behaviours (e.g.  
216 foraging, migration, refuging, nesting, breeding) that drive them to move  
217 against the topographic slope of mountains and hills. Notably, multiple  
218 behaviours can simultaneously shape animal movements and are rarely  
219 mutually exclusive; for instance, foraging and mating often both drive  
220 migration patterns. In a classic example, spawning salmon (*Oncorhynchus*  
221 spp.) annually swim hundreds of kilometers upstream from the ocean,  
222 leading to doubling or tripling of the amount of nitrogen and phosphorous in  
223 certain spawning habitats (Helfield & Naiman, 2001; Holtgrieve & Schindler,  
224 2011; Deacy *et al.*, 2016). Although juveniles eventually migrate (disperse)  
225 back to the ocean, an estimated 85% of marine-derived nutrients from

226carcasses or eggs remain in the food web upstream, supporting both aquatic  
227and terrestrial food webs (Moore & Schindler, 2004). Similarly, insects that  
228exhibit hill-topping behaviour (i.e. aggregating on hills or cliffs to mate;  
229Capinera & Skevington, 2008) could also serve as a subsidy to the regions in  
230which they gather, although this potential nutrient influx has yet to be  
231explicitly quantified.

232       Counter-gradient movement by animal vectors likely occurs in systems  
233where food resources and critical habitat (e.g. for breeding or nesting) are  
234separated by an energy gradient (e.g. uphill) or ecosystem boundary. A  
235prominent example is the vertical movement of deep-diving whales (e.g.  
236*Physeter macrocephalus* and *Balaenoptera physalus*) throughout the water  
237column as they feed in deep ocean layers and then rise to the surface to  
238breathe (known as the ‘whale pump’; Fig. 4; Roman & McCarthy, 2010). In  
239doing so, they move counter-directionally to the downward flux of key  
240nutrients from the surface through the gravitational pull on aggregates and  
241faeces (Roman & McCarthy, 2010). Through this process, cetaceans may  
242transport up to  $2.3 \times 10^4$  metric tons of nitrogen to the surface per year in  
243areas such as the Gulf of Maine – more than all coastal point sources in that  
244region (Roman & McCarthy, 2010). This movement may be particularly  
245important in the open ocean because biological production here depends  
246largely on nutrient dynamics within the photic zone (Longhurst & Harrison,  
2471989). Nutrients from whale waste (i.e. nitrogen) are utilized in this area for  
248growth and photosynthesis by phytoplankton, which are then consumed by

249zooplankton and fish to fuel the biological pump. Without animal movement,  
250these limiting nutrients would be entirely dependent on seasonal upwelling,  
251which is restricted in time and space. Thus, the counter-directional  
252redistribution of nutrients as a result of whale feeding behaviour during  
253migration serves to expand the otherwise seasonal nutrient influx temporally  
254and the locations covered by passive subsidies spatially.

255       Counter-gradient movement of nutrients by active subsidy agents has  
256important implications for ecosystem function over time. This is particularly  
257true when organisms traverse ecosystem boundaries in ways that passive  
258subsidies cannot, enhancing connectivity among systems. For instance, birds  
259can serve as an important conduit of carbon across the marine-terrestrial  
260ecosystem boundary (Anderson & Polis, 1999; Adame *et al.*, 2015;  
261McFadden, Kauffman & Bhomia, 2016; Otero *et al.*, 2018). Classic examples  
262are avian species that forage at sea and nest on islands relatively  
263uninhabited by other organisms (Anderson & Polis, 1999). Unlike the passive  
264exchange by ocean waves of nutrients between marine and terrestrial  
265ecosystems at the island boundary, seabirds consistently forage beyond this  
266boundary, moving ocean nutrients further onto land. On nutrient-limited  
267islands (e.g. desert or mangrove; Fig. 5), this supply of guano to terrestrial  
268plants can also significantly change the nutrient cycles within an island by  
269relieving nutrient limitation (Adame *et al.*, 2015), and the increase in soil  
270nitrogen and phosphorous concentrations has been shown to enhance plant  
271growth and primary productivity (Anderson & Polis, 1999). On desert islands

272in particular, the effect of seabird roosting behaviour can lead to a 3- to 24-  
273fold increase in populations of detritivores, herbivores, and predators, driving  
274productivity and ecosystem-wide dynamics (Anderson & Polis, 1999). In  
275these cases, daily foraging movements create an essential press-like  
276disturbance for these ecosystems.

277       Furthermore, when animal vectors move along specific paths, they  
278have the potential to transport nutrients more rapidly and over larger spatial  
279scales than passive vectors. For example, the migration of sea turtles  
280between foraging (i.e. marine habitat) and nesting grounds (i.e. coastal  
281habitat) represents the long-distance movement of nutrients derived from  
282the feeding grounds to selected coastal ecosystems (Bouchard & Bjørndal,  
2832000), creating a nutrient hotspot *via* both egg content and hatchlings  
284(Bouchard & Bjørndal, 2000). Some species of migrating turtles can transport  
285nutrients over significantly greater distances, at a higher consistency, and  
286over a much shorter time scale than any form of passive subsidy transport  
287(e.g. currents). For example, leatherback turtles (*Dermochelys coriacea*) that  
288feed on jellyfish in the Arctic Circle annually return thousands of kilometers  
289to tropical beaches to nest (Bjørndal, 1997). This nutrient movement  
290supports food webs in the recipient ecosystems, including detrital consumers  
291as well as larger terrestrial predators such as ants, crabs, foxes, and  
292raccoons (Fowler, 1979; Stancyk, 1995). Notably, the spatial distribution of  
293this influx depends on nest-site-selection behaviour. For example,  
294loggerhead turtles (*Caretta caretta*) nest further from dunes than other

295species, and thus the nutrients they transport may contribute less to dune  
296vegetation than to other components of the ecosystem (Witherington, 1986;  
297Bouchard & Bjørndal, 2000). Such targeted transport emphasizes the unique  
298role of behaviour (in this case, habitat selection) in ecosystem subsidies  
299(Subalusky & Post, 2018). Seasonal migrations, like those performed by sea  
300turtles, can create pulsed or cyclic perturbations of nutrient availability  
301within a habitat. Consequently, such counter-gradient movements have the  
302potential to generate important ecosystem impacts that reflect those of  
303passive subsidies (i.e. can be consistent or cyclical in nature), but differ in  
304terms of their pathway.

305

## 306**(2) Agents respond to the nutrient distribution patterns that they** 307**generate**

308       Animal-mediated transport can also be reactive, driven by an  
309organism's behavioural response to its physical environment (e.g.  
310environmental structure, climate, existing resource distribution). Specifically,  
311resource distribution both influences and is influenced by organism  
312movement, and this can lead to positive- or negative-feedback loops in  
313nutrient accumulation (Earl & Zollner, 2017; Subalusky & Post, 2018). This  
314feature of active subsidies is likely most prominent when nutrient influxes  
315affect the resources that agents directly use. Specifically, if a nutrient influx  
316by an animal vector directly affects its own food source (e.g. for a herbivore)  
317or habitat structure (e.g. for a tree-nesting species), this may generate a

318stronger feedback loop compared to situations in which habitat selection is  
319not heavily influenced by nutrient input. However, the direction of the  
320feedback loop (positive or negative) and magnitude of the subsidy input  
321depend on the strength of drivers that influence the behaviour of the active  
322subsidy (see Table 1 for examples).

323       Positive-feedback loops are generated when organisms aggregate in  
324an area of high resource availability and continue to contribute to its  
325productivity through fertilization. One of the best-studied outcomes of this  
326feature is the formation of resource hotspots, in which organisms  
327preferentially revisit areas where they have already deposited nutrients  
328(reviewed in Earl & Zollner, 2017). In these cases, the active subsidy agents  
329can create and/or respond to an influx of resources. This process has been  
330observed when seabird preference for nesting on certain islands has resulted  
331in significantly elevated levels of plant-available nitrogen surrounding areas  
332of concentrated guano deposits (McCauley *et al.*, 2012; Adame *et al.*, 2015;  
333Graham *et al.*, 2018). On nutrient-depleted mangrove islands, for example,  
334seabirds improve the quality of their own habitat through the net influx of  
335nutrient enrichment, rendering the islands more attractive to roosting  
336seabirds and continuing to support – or even increasing suitability for –  
337populations of these active subsidy agents (Adame *et al.*, 2015; Fig. 5).  
338Feedback loops like this increase primary productivity and strengthen  
339autotroph communities (e.g. plants), which not only attracts agent  
340conspecifics, but also heterospecifics. For example, in the African savannah–



341 woodland system, the grassy remnants of former livestock corrals (once  
342 fertilized by livestock dung) often form high-resource glade regions, which in  
343 turn attract ungulate grazers (Augustine, 2003; Muchiru, Western & Reid,  
344 2008). As they forage within this habitat, ungulates fertilize the area with  
345 dung and urine, helping to maintain the productivity of the glade and attract  
346 other species to the area even long after the initial active subsidy agents  
347 have dispersed (Augustine, 2003; Muchiru *et al.*, 2008). These include superb  
348 starlings (*Lamprotornis superbus*), which nest on acacia trees around the  
349 outskirts of glades, taking advantage of the predictable insect abundance  
350 found within glades to provision hatchlings, and thus emphasizing how active  
351 subsidy movement influences both conspecific and heterospecific habitat use  
352 (Rubenstein, 2007, 2016).

353       Conversely, negative-feedback loops occur when organisms actively  
354 avoid certain locations and consequently limit nutrient influx from potential  
355 active subsidies. Most simply, this can occur due to resource depletion, such  
356 as when organisms spread nutrients from hotspots across a broader area  
357 and consequently avoid the initial hotspot locations once the nutrients have  
358 been depleted. On a small spatial scale, this takes place when frugivores  
359 disperse seeds from a tree (a local hotspot), reducing the hotspot and  
360 contributing to a more uniform distribution of resources as seeds from the  
361 fruit are spread from their source (Côtés & Uriarte, 2013). Negative-  
362 feedback loops may also be more complex, driven by biotic interactions as  
363 well as behavioural responses to the environment across wider scales. For

example, on tropical islets in the Central Pacific, invasive coconut trees (*Cocos nucifera*) tend to grow where they benefit from bird-transported nutrients (Young *et al.*, 2010). However, seabirds (e.g. *Sula* spp.) prefer to roost in native trees (*Pisonia grandis* and *Tournefortia argentea*) over these invasive plants (Young *et al.*, 2010). Thus, bird behavioural responses to coconut trees (i.e. repulsion) reduce the initial hotspot by reducing marine-derived subsidies in these locations and triggering nutrient-depletion patterns within the communities on these islands (Young *et al.*, 2010). Negative-feedback loops can also form in existing hotspots when these areas become saturated with nutrients to the point of toxicity ('subsidy overload'; Dutton *et al.*, 2018), as in the case of eutrophication. For example, excessive amounts of nutrient deposition on seabird islands can lead to changes in species composition by killing certain plant species (Hogg & Morton, 1983; Ellis, Fariña & Witman, 2006). In general, since predicting when and how a feedback loop will form depends heavily on the system, considering the net impact of behavioural responses to existing nutrient distribution will require empirical work that is tailored to specific systems.

381

### 382(3) Interactions among agents affect deposition patterns

383       Animal movement is not only affected by abiotic factors, like the  
384physical environment or existing resource distribution, but also by the  
385distribution and movement of other agents (i.e. direct interactions between  
386organisms). These may be conspecific or heterospecific, representing

387competitors, mates, pathogens, prey or predators. Variation in behaviours as  
388a result of other agents, like social attraction or anti-predator responses,  
389thus affects the movement patterns and ecological implications of active  
390subsidies. Furthermore, perturbations of the community composition (e.g.  
391through the introduction of an invasive species or local extinctions) can also  
392affect movement and the derived nutrient deposition patterns, again  
393highlighting how predictions from phenomenological models alone may  
394become inaccurate under changing environmental conditions.

395       Perhaps best studied is how predators influence subsidies through their  
396effect on prey population size and behaviour (Schmitz, Hawlena & Trussell,  
3972010; Strickland *et al.*, 2013). Most obviously, predation can directly affect  
398nutrient deposition within an ecosystem *via* consumptive effects. For  
399example, subsidies from wolf-killed carcasses in Yellowstone National Park  
400have been shown to create temporary hotspots by attracting scavenging  
401species such as ravens (*Corvus corax*), bald eagles (*Haliaeetus*  
402*leucocephalus*), and smaller carnivores (Ripple *et al.*, 2011). But  
403independently of consumptive effects, the ‘landscape of fear’ generated by  
404predator presence also shapes prey habitat preference, food choice, space  
405use and distribution (Laundré, Hernández & Ripple, 2010, Schmitz *et al.*,  
4062010). In turn, evasive behavioural responses to predator activity can affect  
407the distribution of nutrients across a landscape. For instance, after the  
408eradication of Yellowstone wolves (*Canis lupus*) in the early 20th century,  
409relatively uninhibited populations of ungulates nearly doubled the rate of net

410nitrogen availability among grasslands through waste products (Frank,  
4112008). Several years following the reintroduction of grey wolves to the park,  
412there was a notable decrease in net nitrogen availability in these grasslands.  
413This change was likely driven by a shift in ungulate habitat-use patterns,  
414rather than direct removal of these grazers by the reintroduced wolves  
415(Frank, 2008). Thus, in situations where heterospecific interactions play a  
416major role in agent movement, identifying the response (e.g. limited  
417movement or altered patch use) will be critical for predicting how nutrient  
418deposition will change in both space and time.

419       In addition to trophic-level interactions, conspecific interactions can aid  
420in the formation of nutrient hotspots, particularly for species that utilize  
421social information to make decisions about movement within and among  
422habitats (i.e. ‘ecology of information’; Clobert *et al.*, 2009; Schmidt, Dall &  
423Van Gils, 2010). Specifically, social information generates correlations in  
424behaviour and space use of multiple individuals (Gil *et al.*, 2018), thus  
425affecting both the nature and magnitude of active subsidy nutrient  
426distribution across a landscape. For instance, breeding-site selection by  
427many social agents is often based on conspecifics, such as when the  
428presence of nesting marine birds provides cues about local breeding  
429conditions (‘information barrier hypothesis’; Forbes & Kaiser, 1994; Schmidt  
430*et al.*, 2010), deterring individuals from dispersal to a new habitat. In this  
431case, we would hypothesize that the subsequent social aggregation likely  
432drives hotspot formation *via* the accumulation of nutrients in these areas

433(Fig. 1). Central-place foraging behaviour by highly social animals can  
434similarly cause the formation of nutrient hotspots *via* aggregation, especially  
435if these central places remain stable over long periods (Clay *et al.*, 2013). For  
436example, colonial canopy ants (*Azteca trigona*) that continually return to a  
437collective home base following foraging bouts can alter nutrient composition  
438of the leaf litter below their nests *via* a steady stream of nutrient-rich excreta  
439falling on the forest floor (Clay *et al.*, 2013). In these cases, understanding  
440how large, dense and spatially stable conspecific aggregations form is key  
441for predicting the effect of social behaviour on nutrient influxes. Whether  
442aggregating behaviour occurs on a daily (e.g. foraging or anti-predator  
443response) or seasonal (e.g. breeding) basis would then determine the  
444corresponding ecosystem effect (press or pulse), depending on whether the  
445subsidy influx from this movement is consistent or cyclical.

446

#### 447**IV. INDIVIDUAL DIFFERENCES AS A SOURCE OF VARIATION**

448 Although almost entirely overlooked by the ecological subsidy literature,  
449individual variation of within-population space use likely plays a role in  
450generating nutrient-deposition patterns. This variation can arise from a  
451combination of external and internal factors (Nathan *et al.*, 2008). First,  
452individuals might move differently simply because they occupy different  
453habitats with differences in structure and composition. For example,  
454individuals in resource-scarce areas might be forced to move more in search  
455of these resources, resulting in longer transportation distances. Second,

456 differences in movement patterns can arise from variation in individual life  
457 stage, sex, and morphology (Fraser *et al.*, 2001). For instance, winged-morph  
458 insects readily disperse across distances and barriers that de-winged morphs  
459 are physically unable to overcome (Roff, 1986). Furthermore, such  
460 intraspecific differences can occur at multiple scales, reflecting variation  
461 among populations (e.g. along a latitudinal gradient), or differences within a  
462 population. Lastly, a growing body of literature is also linking differential  
463 space-use patterns with variation in life-history traits or behavioural  
464 strategies (Réale *et al.*, 2010; Sih *et al.*, 2012). Personality traits (defined as  
465 within-individual consistency in behavioural responses across time or  
466 contexts) may interact with environmental heterogeneity, resulting in  
467 personality-dependent dispersal and space use that can affect various  
468 ecological processes at the population and community levels (Cote *et al.*  
469 2010a; Spiegel *et al.*, 2017; Villegas-Ríos *et al.*, 2017).

470 Evidence of consistent (phenotype-related) intra-population differences in  
471 dispersal distance, home-range size, habitat selection and other space-use  
472 axes is becoming more common in a wide range of taxa (Duckworth &  
473 Badyaev, 2007; Cote *et al.*, 2010b; Harrison *et al.*, 2014; Spiegel *et al.*,  
474 2015b; Patrick, Pinaud & Weimerskirch, 2017). Understanding these aspects  
475 of behavioural phenotype can provide greater insight into an organism's role  
476 in nutrient distribution, as they can potentially affect the distance, rate, and  
477 amount of nutrients transported. For example, individual wandering  
478 albatross (*Diomedea exulans*) vary in the ratio of their exploration–

479exploitation foraging behaviour (Patrick *et al.*, 2017). Thus, different  
480individuals likely move nutrients between patches at different rates. In this  
481case, one might expect exploiters (i.e. those that spend more time in  
482restricted-area search) to contribute more to local transport and hotspot  
483formation than explorers (i.e. those that spend more time in long-distance  
484travel), while the latter could contribute more to long-range transport and  
485ecosystem connectivity. Similarly, individual brown bears (*Ursus arctos*) vary  
486in the time spent foraging at salmon (*Oncorhynchus* spp.) spawning sites  
487(Deacy *et al.*, 2016). As previously mentioned, salmon are an important  
488source of nutrients in upstream spawning habitats, and their consumers  
489serve as the final link in transporting salmon-derived nutrients from marine  
490and aquatic ecosystems to the terrestrial environment. Variation in salmon  
491consumption by brown bears, as well as in post-consumption space use  
492among individual bears (Leclerc *et al.*, 2016; Hertel *et al.*, 2019), would thus  
493influence the extent to which cross-boundary active subsidy transport  
494occurs.

495 Furthermore, variation in space use can lead to spatial structuring of  
496phenotypes within a population (e.g. Duckworth & Badyaev, 2007; Cote *et*  
497*al.*, 2010a; Spiegel *et al.*, 2017), which could then affect the spatial  
498structuring of nutrient inputs. Examples of such spatial structuring are  
499exemplified by non-native introductions, where individuals found on the  
500invasion front have both morphological and behavioural differences  
501compared to individuals in established areas (e.g. Gruber *et al.* 2017). In

502these cases, larger and bolder individuals commonly found at the dispersal  
503front would be expected to deposit greater amounts of subsidies at further  
504distances. Scenarios like this suggest that incorporating intraspecific  
505differences in space use can help provide a better understanding of how the  
506same population of animal vectors may simultaneously generate differing  
507patterns of nutrient deposition within and among habitats.

508

## 509**V. CONSERVATION IMPLICATIONS**

510       There is increasing recognition that the materials exchanged *via*  
511animal vector links should be incorporated into management decisions to  
512maintain resilience and ensure future ecosystem function (Lundberg &  
513Moberg, 2003). Previous work has highlighted some of the major implications  
514of removing active subsidy agents in altered ecosystems (Subalusky & Post,  
5152018). Among these, human-induced rapid environmental change (HIREC;  
516Sih, Ferrari & Harris, 2011) can cause population reductions or even local  
517extinctions, resulting in fewer active subsidy agents (Barnosky *et al.*, 2011;  
518Dirzo *et al.*, 2014; Earl & Zollner, 2017; Subalusky & Post, 2018). For  
519example, the movement of nutrients from sea to land *via* seabirds and  
520anadromous fish has been reduced by 96% due to the loss of these species  
521(Doughty *et al.*, 2016). The reduction of animal vector populations and  
522subsequent active subsidy influxes also has a variety of indirect ecological  
523effects. For instance, moth migration altered by changes in large-scale  
524weather patterns affects the movement of Brazilian free-tailed bats



525(*Tadarida brasiliensis*), since these bats rely heavily on this seasonal  
526resource to gain fat for their own autumn migrations (Krauel *et al.*, 2018).

527       Although the number of applied models incorporating organism  
528movement is increasing (Holyoak *et al.*, 2008; Grüss *et al.*, 2011),  
529understanding how behavioural responses change in altered environments  
530can lend greater predictive power to changes in active subsidy nutrient  
531distribution than considering movement patterns alone (Fraser *et al.*, 2018).  
532For example, HIREC could reduce the extent of active subsidy agent  
533movement by increasing fragmentation (i.e. creating physical barriers to  
534movement) or by generating a plastic change in behavioural response (e.g.  
535alterations in habitat use or home-range size). In addition, because many  
536large species tend to travel further, they may exhibit higher sensitivity in  
537behavioural responses to anthropogenic change, altering their movement  
538more significantly and thus inhibiting their ability to act as active subsidies  
539(Tucker *et al.*, 2018). Moreover, HIREC can affect the phenotypic composition  
540within populations, through selection of certain life-history traits, behavioural  
541types or morphologies (e.g. Cooke *et al.*, 2007; Biro & Post, 2008; Parsons,  
542Morrison & Slater, 2010). For example, human harvesting selects against  
543phenotypes with large size, fast growth rates, and high activity and boldness  
544(Biro & Post, 2008; Huntingford, Mesquita & Kadri, 2013). At both local and  
545broader scales, removal of more mobile phenotypes within certain species  
546would limit nutrient distribution by active subsidy agents. Conversely, for  
547recently introduced species, selection may favour bolder individuals that

548 might achieve greater dispersal distances (e.g. Duckworth & Badyaev, 2007)  
549 and therefore transport nutrients further into novel environments. A third  
550 scenario is also possible, by which more behaviourally plastic species or  
551 individuals are best able to adapt to HIREC by calibrating their behaviour to  
552 the current circumstances (e.g. Crowley *et al.*, 2019). For example, recent  
553 modelling of Burmese python (*Python molurus bivittatus*) behaviour suggests  
554 that species able to exhibit adaptively flexible dispersal behaviour are most  
555 successful in population expansion (Mutascio *et al.*, 2017).

556       In general, one critical step for managers and policy-makers will be to  
557 recognize the limitations of spatially restricted forms of management to  
558 enact more proactive conservation measures for mobile animal vectors. For  
559 such animals in the pelagic environment, including sharks and whales,  
560 alternative options could include dynamic (i.e. mobile) or seasonal marine  
561 protected areas to cover key locations in a given species' range (Game *et al.*,  
562 2009). In terrestrial habitats, movement corridors could protect large  
563 migratory or far-ranging species (Silveira *et al.*, 2014; Spiegel *et al.*, 2015a;  
564 Belote *et al.*, 2016) such as raptors and insects. Regardless of management  
565 form, given the alarming evidence of reduction in biomass and movement  
566 ranges of various species exposed to increasing levels of anthropogenic  
567 change, we suggest prioritizing conservation approaches that facilitate the  
568 maintenance of active subsidy behaviour across taxa.

569

## 570 VI. CONCLUSIONS

571(1) The role of animal behaviour in the flow of resources has been  
572underappreciated. Here we suggest that consideration of the behaviours that  
573drive animal movement patterns can provide a better understanding of  
574ecological processes.

575(2) Animal behaviours operate at various spatial and temporal scales, and  
576can generate effects that are quantitatively similar to passive subsidies in  
577nature (i.e. press and pulse perturbation) and, in some cases, can be equal  
578or greater in magnitude.

579(3) Active subsidy transport processes differ qualitatively from those of  
580passive subsidies in that they are behaviourally driven. Three features of  
581animal behaviour highlight how active subsidy behaviours lead to nutrient-  
582distribution patterns unachievable by passive subsidies: (1) animals can  
583move against abiotic gradients; (2) animals respond to the distribution of  
584nutrients they help form (e.g. positive-feedback loops and the formation of  
585nutrient hotspots); (3) animals also interact with other heterospecific and  
586conspecific active subsidy agents, altering their nutrient-deposition patterns.  
587 (4) Individuals within populations of animal vectors often exhibit consistent  
588behavioural differences, which can affect their movement patterns and  
589generate variation in nutrient transport. While widely acknowledged in the  
590context of movement and behaviour, these recent insights have been under-  
591investigated in the context of active subsidies.

592(5) Lastly, it is important to consider animal behaviour when predicting  
593ecosystem-wide responses to future environmental alteration. Specifically,

594human-induced rapid environmental change (HIREC) can interfere with the  
595proposed behavioural features by altering not only the number of acting  
596agents or the community composition, but also their space use and  
597consequent nutrient-deposition patterns. Thus, integrating animal behaviour  
598into predictive models of nutrient fluxes will help to assess more accurately  
599the mechanisms that drive variation in nutrient cycling and how these will be  
600disturbed in the future.

601

602

## 603**VII. ACKNOWLEDGEMENTS**

604We thank Andy Sih, Jenn Rehage and three anonymous reviewers for their  
605helpful comments. We would also like to recognize Eric Van Cleave and  
606Matthew Savoca, who provided valuable insights and reference suggestions.  
607Finally, A.G.M. thanks Damien Caillaud and Nann Fangue for their guidance  
608and mentorship.

609

## 610**VIII. REFERENCES**

611

612ADAME, M.F., FRY, B., GAMBOA, J.N. & HERRERA-SILVEIRA, J.A. (2015) Nutrient  
613 subsidies delivered by seabirds to mangrove islands. *Marine Ecology*  
614 *Progress Series* **525**, 15–24.

615ALLGEIER, J.E., BURKEPILE, D.E. & LAYMAN, C.A. (2017) Animal pee in the sea:  
616 consumer-mediated nutrient dynamics in the world's changing oceans.  
617 *Global Change Biology* **23**, 2166–2178.

618ANDERSON, W.B. & POLIS, G.A. (1999) Nutrient fluxes from water to land:  
619 seabirds affect plant nutrient status on Gulf of California islands.  
620 *Oecologia* **118**, 324–332.

- 621AUGUSTINE, D.J. (2003) Long-term, livestock-mediated redistribution of  
 622 nitrogen and phosphorus in an East African savanna. *Journal of Applied*  
 623 *Ecology* **40**, 137-149.
- 624BARNOSKY, A.D., MATZKE, N., TOMIYA, S., WOGAN, G.O., SWARTZ, B., QUENTAL, T.B.,  
 625 MARSHALL, C., MCGUIRE, J.L., LINDSEY, E.L., MAGUIRE, K.C. & MERSEY, B. (2011)  
 626 Has the Earth's sixth mass extinction already arrived? *Nature* **471**, 51-  
 627 57.
- 628BARNOSKY, A.D., HADLY, E.A., BASCOMPTE, J., BERLOW, E.L., BROWN, J.H., FORTELIUS,  
 629 M., GETZ, W.M., HARTE, J., HASTINGS, A., MARQUET, P.A., MARTINEZ, N.D.,  
 630 MOOERS, A., ROOPNARINE, P., VERMEIJ, G., WILLIAMS, J.W., ET AL. (2012)  
 631 Approaching a state shift in Earth's biosphere. *Nature* **486**, 52-58.
- 632BAUER, S. & HOYE, B.J. (2014) Migratory animals couple biodiversity and  
 633 ecosystem functioning worldwide. *Science* **344**, 1242552.
- 634BELOTE, R.T., DIETZ, M.S., MCRAE, B.H., THEOBALD, D.M., MCCLURE, M.L., IRWIN,  
 635 G.H., MCKINLEY, P.S., GAGE, J.A. & APLET, G.H. (2016) Identifying corridors  
 636 among large protected areas in the United States. *PLoS ONE* **11**.
- 637BENDER, T.J., CASE, E.A. & GILPIN, M.E. (1984) Perturbation experiments in  
 638 community ecology: theory and practice. *Ecology* **65**, 1-13.
- 639BIRD, M.I., BOOBYER, E.M., BRYANT, C., LEWIS, H.A., PAZ, V. & STEPHENS, W.E.  
 640 (2007) A long record of environmental change from bat guano deposits  
 641 in Makangit Cave, Palawan, Philippines. *Earth and Environmental Science*  
 642 *Transactions of the Royal Society of Edinburgh* **98**, 59-69.
- 643BIRO, P.A. & POST, J.R. (2008) Rapid depletion of genotypes with fast growth  
 644 and bold personality traits from harvested fish population. *Proceedings*  
 645 *of the Natinoal Academy of Sciences* **105**, 2919-2922.
- 646BJØRNDAL, K.A. (1997) Foraging ecology and nutrition of sea turtles. In *The*  
 647 *biology of sea turtles* (eds P.L. LUTZ & J.A. MUSICK), pp. 199-232. CRC  
 648 Press, Boca Raton, FL.
- 649BOTH, C., BOUWHUIS, S., LESSELLS, C.M. & VISSER, M.E. (2006) Climate change  
 650 and population declines in a long-distance migratory bird. *Nature* **441**,  
 651 81.
- 652BOUCHARD, S.S. & BJØRNDAL, K.A. (2000) Sea turtles as biological transporters  
 653 of nutrients and energy from marine to terrestrial ecosystems. *Ecology*  
 654 **81**, 2305-2313.

- 655CAPINERA, J.L. & SKEVINGTON, J.H. (2008) Hilltopping. In *Encyclopedia of*  
656 *Entomology, Second Edition*. Kluwer Academic Publisher.
- 657CAPPS, K.A. & FLECKER, A.S. (2013) Invasive fishes generate biogeochemical  
658 hotspots in a nutrient-limited system. *PLoS ONE* **8**, e54093.
- 659CHILDRESS, E.S. & MCINTYRE, P.B. (2015) Multiple nutrient subsidy pathways  
660 from a spawning migration of iteroparous fish. *Freshwater Biology* **60**,  
661 490–499.
- 662CLAY, N.A., LUCAS, J., KASPARI, M. & KAY, A.D. (2013) Manna from heaven:  
663 refuse from an arboreal ant links aboveground and belowground  
664 processes in a lowland tropical forest. *Ecosphere* **4**, 1–15.
- 665CLEVELAND, C.C., HOULTON, B.Z., SMITH, W.K., MARKLEIN, A.R., REED, S.C., PARTON,  
666 W., DEL GROSSO, S.J. & RUNNING, S.W. (2013) Patterns of new versus  
667 recycled primary production in the terrestrial biosphere. *Proceedings of*  
668 *the National Academy of Sciences* **110**, 12733–12737.
- 669CLOBERT, J., LE GALLIARD, J.-F., COTE, J., MEYLAN, S. & MASSOT, M. (2009) Informed  
670 dispersal, heterogeneity in animal dispersal syndromes and the  
671 dynamics of spatially structured populations. *Ecology Letters* **12**, 197–  
672 209.
- 673COOK, P.J. & MCELHINNY, M.W. (1979) A reevaluation of the spatial and  
674 temporal distribution of sedimentary phosphate deposits in the light of  
675 plate tectonics. *Economic Geology* **74**, 315–330.
- 676COOKE, S.J., SUSKI, C.D., OSTRAND, K.G., WAHL, D.H. & PHILIPP, D.P. (2007)  
677 Physiological and behavioral consequences of long-term artificial  
678 selection for vulnerability to recreational angling in a teleost fish.  
679 *Physiological and Biochemical Zoology* **80**, 480–490.
- 680CÔRTEZ, M.C. & URIARTE, M. (2013) Integrating frugivory and animal  
681 movement: a review of the evidence and implications for scaling seed  
682 dispersal. *Biological Reviews* **88**, 255–272.
- 683COTE, J., CLOBERT, J., BRODIN, T., FOGARTY, S. & SIH, A. (2010a) Personality-  
684 dependent dispersal: characterization, ontogeny and consequences for  
685 spatially structured populations. *Philosophical Transactions of the Royal*  
686 *Society B: Biological Sciences* **365**, 4065–4076.
- 687COTE, J., FOGARTY, S., WEINERSMITH, K., BRODIN, T. & SIH, A. (2010b) Personality  
688 traits and dispersal tendency in the invasive mosquitofish (*Gambusia*  
689 *affinis*). *Proceedings of the Royal Society of London B: Biological*  
690 *Sciences* **277**, 1571–1579.

691 CROWLEY, P.H., TRIMMER, P.C., SPIEGEL, O., EHLMAN, S.M., CUELLO, W.S. & SIH, A.  
692 (2019) Predicting habitat choice after rapid environmental change. *The*  
693 *American Naturalist* **193**, 619–632.

694 DEACY, W., LEACOCK, W., ARMSTRONG, J.B. & STANFORD, J.A. (2016) Kodiak brown  
695 bears surf the salmon red wave: direct evidence from GPS collared  
696 individuals. *Ecology* **97**, 1091–1098.

697 DIRZO, R., YOUNG, H.S., GALETTI, M., CEBALLOS, G., ISAAC, N.J.B. & COLLEN, B.  
698 (2014) Defaunation in the anthropocene. *Science* **401**, 401–406.

699 DOUGHTY, C.E., ROMAN, J., FAURBY, S., WOLF, A., HAQUE, A., BAKKER, E.S., MALHI,  
700 Y., DUNNING, J.B. & SVENNING, J.-C. (2016) Global nutrient transport in a  
701 world of giants. *Proceedings of the National Academy of Sciences* **113**,  
702 868–873.

703 DREYER, J., TOWNSEND, P.A., HOOK, J.C., HOEKMAN, D., VANDER ZANDEN, M.J. &  
704 GRATTON, C. (2015) Quantifying aquatic insect deposition from lake to  
705 land. *Ecology* **96**, 499–509.

706 DUCKWORTH, R.A. & BADYAEV, A. V (2007) Coupling of dispersal and aggression  
707 facilitates the rapid range expansion of a passerine bird. *Proceedings of*  
708 *the National Academy of Sciences* **104**, 15017–15022.

709 DUTTON, C.L., SUBALUSKY, A.L., HAMILTON, S.K., ROSI, E.J. & POST, D.M. (2018)  
710 Organic matter loading by hippopotami causes subsidy overload  
711 resulting in downstream hypoxia and fish kills. *Nature Communications*  
712 **9**, 1951.

713 EARL, J.E. & ZOLLNER, P.A. (2014) Effects of animal movement strategies and  
714 costs on the distribution of active subsidies across simple landscapes.  
715 *Ecological Modeling* **283**, 45–52.

716 EARL, J.E. & ZOLLNER, P.A. (2017) Advancing research on animal-transported  
717 subsidies by integrating animal movement and ecosystem modeling.  
718 *Journal of Animal Ecology* **86**, 987–997.

719 ELLIS, J.C., FARIÑA, J.M. & WITMAN, J.D. (2006) Nutrient transfer from sea to  
720 land: the case of gulls and cormorants in the Gulf of Maine. *Journal of*  
721 *Animal Ecology* **75**, 565–574.

722 ELSE, J.J., BRACKEN, M.E., CLELAND, E.E., GRUNER, D.S., HARPOLE, W.S.,  
723 HILLEBRAND, H., NGAI, J.T., SEABLOOM, E.W., SHURIN, J.B. & SMITH, J.E. (2007)  
724 Global analysis of nitrogen and phosphorus limitation of primary  
725 producers in freshwater, marine and terrestrial ecosystems. *Ecology*  
726 *Letters* **10**, 1135–1142.

- 727FAGAN, W.F., LUTSCHER, F. & SCHNEIDER, K. (2007) Population and community  
728 consequences of spatial subsidies derived from central-place foraging.  
729 *The American Naturalist* **170**, 902–915.
- 730FENOLIO, D.B., GRAENING, G.O., COLLIER, B.A. & STOUT, J.F. (2006) Coprophagy in  
731 a cave-adapted salamander; the importance of bat guano examined  
732 through nutritional and stable isotope analyses. *Proceedings of the Royal*  
733 *Society B: Biological Sciences* **273**, 439–443.
- 734FERREIRA, R.L. & MARTINS, R.P. (1999) Trophic structure and natural history of  
735 bat guano invertebrate communities, with special reference to Brazilian  
736 caves. *Tropical Zoology* **12**, 231–252.
- 737FORBES, L. & KAISER, G.W. (1994) Habitat choice in breeding seabirds: when to  
738 cross the information barrier. *Oikos* **70**, 377–384.
- 739FOWLER, L.E. (1979) Hatching success and nest predation in the green sea  
740 turtle, *Chelonia Mydas*, at Tortuguero, Costa Rica. *Ecology* **60**, 946–955.
- 741FRANK, D.A. (2008) Evidence for top predator control of a grazing ecosystem.  
742 *Oikos* **117**, 1718–1724.
- 743FRASER, D.F., GILLIAM, J.F., DALEY, M.J., LE, A.N. & SKALSKI, G.T. (2001) Explaining  
744 leptokurtic movement distributions: intrapopulation variation in boldness  
745 and exploration. *The American Naturalist* **158**, 124–135.
- 746FRASER, K.C., DAVIES, K.T.A., DAVY, C.M., FORD, A.T., FLOCKHART, D.T.T. &  
747 MARTINS, E.G. (2018) Tracking the conservation promise of movement  
748 ecology. *Frontiers in Ecology and Evolution* **6**, 1–7.
- 749GAME, E., GRANTHAM, H., HOBDAI, A.J., PRESSEY, R.I., LOMBARD, A.T., BECKLEY, L.E.,  
750 GJERDE, K., BUSTAMANTE, R., POSSINGHAM, H. & RICHARDSON, A.J. (2009)  
751 Pelagic protected areas: the missing dimension in ocean conservation.  
752 *Trends in Ecology and Evolution* **24**, 360–369.
- 753GENDE, S.M., EDWARDS, R.T., WILLSON, M.F. & WIPFLI, M.S. (2002) Pacific  
754 salmon in aquatic and terrestrial ecosystems. *American Institute of*  
755 *Biological Sciences* **52**, 917–928.
- 756GIL, M., HEIN, A., SPIEGEL, O., BASKETT, M.L. & SIH, A. (2018) Social information  
757 links individual behavior to population and community dynamics. *Trends*  
758 *in Ecology and Evolution* **33**, 535–548.
- 759GRAHAM, N.A.J., WILSON, S.K., CARR, P., HOEY, A.S., JENNINGS, S. & McNeil, M.A.  
760 (2018) Seabirds enhance coral reef productivity and functioning in the  
761 absence of invasive rats. *Nature* **559**, 250–253.



- 762 GRUBER, J., BROWN, G., WHITING, M.J. & SHINE, R. (2017) Is the behavioural  
763 divergence between range-core and range-edge populations of cane  
764 toads (*Rhinella marina*) due to evolutionary change or developmental  
765 plasticity? *Royal Society Open Science* **4**, 1–9.
- 766 GRÜSS, A., KAPLAN, D.M., GUÉNETTE, S., ROBERTS, C.M. & BOTSFORD, L.W. (2011)  
767 Consequences of adult and juvenile movement for marine protected  
768 areas. *Biological Conservation* **144**, 692–702.
- 769 HAMBURG, S.P. & LIN, T.-C. (1998) Throughfall chemistry of an ecotonal forest  
770 on the edge of the Great Plains. *Canadian Journal of Forest Research* **28**,  
771 1456–1463.
- 772 HARRISON, P.M., GUTOWSKY, L.F.G., MARTINS, E.G., PATTERSON, D.A., COOKE, S.J. &  
773 POWER, M. (2014) Personality-dependent spatial ecology occurs  
774 independently from dispersal in wild burbot (*Lota lota*). *Behavioral*  
775 *Ecology* **26**, 483–492.
- 776 HATA, H., NISHIHARA, M. & KAMURA, S. (2002) Effects of habitat-conditioning by  
777 the damselfish *Stegastes nigricans* (Lacepède) on the community  
778 structure of benthic algae. *Journal of Experimental Marine Biology and*  
779 *Ecology* **280**, 95–116.
- 780 HELFIELD, J.M. & NAIMAN, R.J. (2001) Effects of salmon-derived nitrogen on  
781 riparian forest growth and implications for stream productivity. *Ecology*  
782 **82**, 2403–2409.
- 783 HELFIELD, J.M. & NAIMAN, R.J. (2006) Keystone interactions: salmon and bear in  
784 riparian forests of Alaska. *Ecosystems* **9**, 167–180.
- 785 HERTEL, A., LECLERC, M., WARREN, D., PELLETIER, F., ZEDROSSER, A. & MUELLER, T.  
786 (2019) Don't poke the bear: using tracking data to quantify behavioural  
787 syndromes in elusive wildlife. *Animal Behaviour* **147**, 91–104.
- 788 HOGG, E.H. & MORTON, J.K. (1983) The effects of nesting gulls on the  
789 vegetation and soil of islands in the Great Lakes. *Canadian Journal of*  
790 *Botany* **61**, 3240–3254.
- 791 HOLDO, R.M., HOLT, R.D., COUGHENOUR, M.B. & RITCHIE, M.E. (2007) Plant  
792 productivity and soil nitrogen as a function of grazing, migration and fire  
793 in an African savanna. *Journal of Ecology* **95**, 115–128.
- 794 HOLTGRIEVE, G.W. & SCHINDLER, D.E. (2011) Marine-derived nutrients,  
795 bioturbation, and ecosystem metabolism: reconsidering the role of  
796 salmon in streams. *Ecology* **92**, 373–385.

797 HOLYOAK, M., CASAGRANDE, R., NATHAN, R., REVILLA, E. & SPIEGEL, O. (2008)  
798 Trends and missing parts in the study of movement ecology.  
799 *Proceedings of the National Academy of Sciences* **105**, 19060–19065.

800 HOULTON, B. & MORFORD, S. (2015) A new synthesis for terrestrial nitrogen  
801 inputs. *Soil* **1**, 381–397.

802 HU, K., HORVITZ, N., CLARK, S., REYNOLDS, D., SAPIR, N. & CHAPMAN, J.G. (2017)  
803 Mass seasonal bioflows of high-flying insect migrants. *Science* **354**,  
804 1584–1586.

805 HUNTINGFORD, B., MESQUITA, F. & KADRI, S. (2013) Personality variation in  
806 cultured fish: implications for production and welfare. In *Animal*  
807 *personalities: Behavior, physiology and evolution* (eds C. CARERE & D.  
808 MAESTRIPIERI), pp. 414–440. University of Chicago Press, Chicago.

809 JELTSCH, F., BONTE, D., PE'ER, G., REINEKING, B., LEIMGRUBER, P., BALKENHOL, N.,  
810 SCHRÖDER, B., BUCHMANN, C.M., MUELLER, T., BLAUM, N., ZURELL, D. &  
811 THORSTEN, K.B. (2013) Integrating movement ecology with biodiversity  
812 research. *Movement Ecology* **1**, 1–13.

813 KRAUEL, J.J., RATCLIFFE, J.M., WESTBROOK, J.K. & MCCracken, G.F. (2018) Brazilian  
814 free-tailed bats (*Tadarida brasiliensis*) adjust foraging behaviour in  
815 response to migratory moths. *Canadian Journal of Zoology* **96**, 513–520.

816 KUNZ, T.H., BRAUN DE TORREZ, E., BAUER, D., LOBOVA, T. & FLEMING, T.H. (2011)  
817 Ecosystem services provided by bats. *Annals of the New York Academy*  
818 *of Sciences* **1223**, 1–38.

819 LAUNDRÉ, J.W., HERNANDEZ, L. & RIPPLE, W.J. (2010) The landscape of fear:  
820 ecological implications of being afraid. *The Open Ecology Journal* **3**, 1–7.

821 LEBAUER, D.S. & TRESEDER, K.K. (2008) Nitrogen limitation of net primary  
822 productivity in terrestrial ecosystems is globally distributed. *Ecology* **89**,  
823 371–379.

824 LECLERC, M., VANDER WAL, E., ZEDROSSER, A., SWENSON, J.E., KINDBERG, J. &  
825 PELLETIER, F. (2016) Quantifying consistent individual differences in  
826 habitat selection. *Oecologia* **180**, 697–705.

827 LONGHURST, A.R. & HARRISON, W.G. (1989) The biological pump: profiles of  
828 plankton production and consumption in the upper ocean. *Progress in*  
829 *Oceanography* **22**, 47–123.

- 830 LUNDBERG, J. & MOBERG, F. (2003) Mobile link organisms and ecosystem  
831 functioning: implications for ecosystem resilience and management.  
832 *Ecosystems* **6**, 0087–0098.
- 833 MARCARELLI, A.M., BAXTER, C. V., MINEAU, M.M. & HALL, R.O. (2011) Quantity and  
834 quality: unifying food web and ecosystem perspectives on the role of  
835 resource subsidies in freshwaters. *Ecology* **92**, 1215–1225.
- 836 MCCANN, N.P., ZOLLNER, P.A. & GILBERT, J.H. (2017) Temporal scaling in analysis  
837 of animal activity. *Ecography* **40**, 1436–1444.
- 838 MCCAULEY, D.J., DESALLES, P.A., YOUNG, H.S., DUNBAR, R.B., DIRZO, R., MILLS, M.M.  
839 & MICHELI, F. (2012) From wing to wing: the persistence of long ecological  
840 interaction chains in less-disturbed ecosystems. *Scientific Reports* **2**, 1–9.
- 841 MCFADDEN, T.N., KAUFFMAN, J.B. & BHOMIA, R.K. (2016) Effects of nesting  
842 waterbirds on nutrient levels in mangroves, Gulf of Fonseca, Honduras.  
843 *Wetlands Ecology and Management* **24**, 217–229.
- 844 MCINTYRE, P.B., FLECKER, A.S., VANNI, M.J., HOOD, J.M., TAYLOR, B.W. & THOMAS,  
845 S.A. (2008) Fish distributions and nutrient cycling in streams: can fish  
846 create biogeochemical hotspots. *Ecology* **89**, 2335–2346.
- 847 MCNAUGHTON, S.J., BANYIKWA, F.F. & MCNAUGHTON, M.M. (1997) Promotion of the  
848 cycling of diet-enhancing nutrients by African grazers. *Science* **278**,  
849 1798–1800.
- 850 MOORE, J.W. & SCHINDLER, D.E. (2004) Nutrient export from freshwater  
851 ecosystems by anadromous sockeye salmon (*Oncorhynchus nerka*).  
852 *Canadian Journal of Fisheries and Aquatic Sciences* **61**, 1582–1589.
- 853 MORFORD, S.L., HOULTON, B.Z. & DAHLGREN, R.A. (2016) Geochemical and  
854 tectonic uplift controls on rock nitrogen inputs across terrestrial  
855 ecosystems. *Global Biogeochemical Cycles* **30**, 339–349.
- 856 MUCHIRU, A.N., WESTERN, D.J. & REID, R.S. (2008) The role of abandoned  
857 pastoral settlements in the dynamics of African large herbivore  
858 communities. *Journal of Arid Environments* **72**, 940–952.
- 859 MUTASCIO, H.E., PITTMAN, S.E. & ZOLLNER, P.A. (2017) Investigating movement  
860 behavior of invasive Burmese pythons on a shy–bold continuum using  
861 individual-based modeling. *Perspectives in Ecology and Conservation* **15**,  
862 25–31.
- 863 NATHAN, R., GETZ, W., REVILLA, E., HOLYOAK, M., KADMON, R., SALTZ, D., & SMOUSE,  
864 P. E., R, W.M. (2008) A movement ecology paradigm for unifying

865 organismal movement research. *Proceedings of the National Academy of*  
866 *Sciences* **105**, 19052–19059.

867 NELSON, J.A., STALLINGS, C.D., LANDING, W.M. & CHANTON, J. (2013) Biomass  
868 transfer subsidizes nitrogen to offshore food webs. *Ecosystems* **16**,  
869 1130–1138.

870 OTERO, X.L., DE LA PEÑA-LASTRA, S., PÉREZ-ALBERTI, A., FERREIRA, T.O. & HUERTA-  
871 DIAZ, M.A. (2018) Seabird colonies as important global drivers in the  
872 nitrogen and phosphorus cycles. *Nature Communications* **9**, 246.

873 PARSONS, D.M., MORRISON, M.A. & SLATER, M.J. (2010) Responses to marine  
874 reserves: decreased dispersion of the sparid *Pagrus auratus* (snapper).  
875 *Biological Conservation* **143**, 2039–2048.

876 PATRICK, S.C., PINAUD, D. & WEIMERSKIRCH, H. (2017) Boldness predicts an  
877 individual's position along an exploration–exploitation foraging trade-off.  
878 *Journal of Animal Ecology* **86**, 1257–1268.

879 PETERJOHN, W.T. & CORRELL, D.L. (1984) Nutrient dynamics in an agricultural  
880 watershed: observations on the role of riparian forest. *Ecology* **65**, 1466–  
881 1475.

882 POLIS, G.A., ANDERSON, W.B. & HOLT, R.D. (1997) Toward an integration of  
883 landscape and food web ecology: the dynamics of spatially subsidized  
884 food webs. *Annual Review of Ecology and Systematics* **28**, 289–316.

885 POLUNIN, N.V.. & KOIKE, I. (1987) Temporal focusing of nitrogen release by a  
886 periodically feeding herbivorous reef fish. *Journal of Experimental Marine*  
887 *Biology and Ecology* **111**, 285–296.

888 POULSON, T.L. & LAVOIE, K.H. (2000) The trophic basis of subsurface  
889 ecosystems. In *Ecosystems of the World: Subterranean Ecosystems* (eds  
890 H. WILKENS, D.C. CULVER & W.F. HUMPHREYS), pp. 231–249. Elsevier  
891 Science.

892 RÉALE, D., GARANT, D., HUMPHRIES, M.M., BERGERON, P., CAREAU, V. & MONTIGLIO,  
893 P.O. (2010) Personality and the emergence of the pace-of-life syndrome  
894 concept at the population level. *Philosophical Transactions of the Royal*  
895 *Society Biological Sciences* **365**, 4051–4063.

896 RIPPLE, W.J., WIRSING, A.J., BESCHTA, R.L. & BUSKIRK, S.W. (2011) Can restoring  
897 wolves aid in lynx recovery? *Wildlife Society Bulletin* **35**, 514–518.

898 ROFF, D.A. (1986) The evolution of wing dimorphism in insects. *Evolution* **40**,  
899 1009–1020.

- 900ROMAN, J. & MCCARTHY, J.J. (2010) The whale pump: marine mammals enhance  
901 primary productivity in a coastal basin. *PLoS ONE* **5**, e13255.
- 902RUBENSTEIN, D. (2016) Superb starlings: cooperation and conflict in an  
903 unpredictable environment. In *Cooperative breeding in vertebrates:  
904 Studies of ecology, evolution, and behavior* (eds W.D. KOENIG & J.L.  
905 DICKINSON), pp. 181–196. Cambridge University Press.
- 906RUBENSTEIN, D.R. (2007) Territory quality drives intraspecific patterns of  
907 extrapair paternity. *Behavioral Ecology* **18**, 1058–1064.
- 908SATO, T., WATANABE, K., KANAIWA, M., NIIZUMA, Y., HARADA, Y. & LAFFERTY, K.D.  
909 (2011) Nematomorph parasites drive energy flow through a riparian  
910 ecosystem. *Ecology* **92**, 201–207.
- 911SCHMIDT, K.A., DALL, S.R.X. & VAN GILS, J.A. (2010) The ecology of information:  
912 an overview on the ecological significance of making informed decisions.  
913 *Oikos* **119**, 304–316.
- 914SCHMITZ, O.J., HAWLENA, D. & TRUSSELL, G.C. (2010) Predator control of  
915 ecosystem nutrient dynamics. *Ecology Letters* **13**, 1199–1209.
- 916SEAGLE, S.W. (2003) Can ungulates foraging in a multiple-use landscape alter  
917 forest nitrogen budgets? *Oikos* **103**, 230–234.
- 918SIH, A., COTE, J., EVANS, M., FOGARTY, S. & PRUITT, J. (2012) Ecological  
919 implications of behavioural syndromes. *Ecology Letters* **15**, 278–289.
- 920SIH, A., FERRARI, M.C.O. & HARRIS, D.J. (2011) Evolution and behavioural  
921 responses to human-induced rapid environmental change. *Evolutionary  
922 Applications* **4**, 367–387.
- 923SILVEIRA, L., SOLLMANN, R., JÁCOMO, A.T.A., DINIZ FILHO, J.A.F. & TÔRRES, N.M.  
924 (2014) The potential for large-scale wildlife corridors between protected  
925 areas in Brazil using the jaguar as a model species. *Landscape Ecology*  
926 **29**, 1213–1223.
- 927SOWLES, J. (2001) Nitrogen in the Gulf of Maine: sources, susceptibility, and  
928 trends. *NOAA/UNH Cooperative Institute for Coastal and Estuarine  
929 Environmental Technology, Gulf of Maine Council on the Marine  
930 Environment, & NOAA Ocean Service*.
- 931SPIEGEL, O., HAREL, R., CENTENO-CUADROS, A., HATZOFÉ, O., GETZ, W.M. & NATHAN,  
932 R. (2015a) Moving beyond curve fitting: using complementary data to  
933 assess alternative explanations for long movements of three vulture  
934 species. *The American Naturalist* **185**, 44–54.

- 935 SPIEGEL, O., LEU, S.T., BULL, C.M. & SIH, A. (2017) What's your move?  
 936 Movement as a link between personality and spatial dynamics in animal  
 937 populations. *Ecology Letters* **20**, 3–18.
- 938 SPIEGEL, O., LEU, S.T., SIH, A., GODFREY, S.S. & BULL, C.M. (2015b) When the  
 939 going gets tough: behavioural type-dependent space use in the sleepy  
 940 lizard changes as the season dries. *Proceedings of the Royal Society of*  
 941 *London B: Biological Sciences* **282**, 20151768.
- 942 STANCYK, S.E. (1995) Non-human predators of sea turtles and their control. In  
 943 *Biology and conservation of sea turtles* (ed K.A. BJORN DAL), pp. 139–152.  
 944 Smithsonian Institution Press, Washington, D.C.
- 945 STRICKLAND, M.S., HAWLENA, D., REESE, A., BRADFORD, M.A. & SCHMITZ, O.J. (2013)  
 946 Trophic cascade alters ecosystem carbon exchange. *Proceedings of the*  
 947 *National Academy of Sciences* **110**, 11035–11038.
- 948 SUBALUSKY, A.L., DUTTON, C.L., ROSI-MARSHALL, E.J. & POST, D.M. (2015) The  
 949 hippopotamus conveyor belt: vectors of carbon and nutrients from  
 950 terrestrial grasslands to aquatic systems in sub-Saharan Africa.  
 951 *Freshwater Biology* **60**, 512–525.
- 952 SUBALUSKY, A.L., DUTTON, C.L., ROSI, E.J. & POST, D.M. (2017) Annual mass  
 953 drownings of the Serengeti wildebeest migration influence nutrient  
 954 cycling and storage in the Mara River. *Proceedings of the National*  
 955 *Academy of Sciences* **114**, 7647–7652.
- 956 SUBALUSKY, A.L. & POST, D.M. (2019) Context dependency of animal resource  
 957 subsidies. *Biological Reviews* **94**, 517–538.
- 958 TOWNSEND, D.W. (1998) Sources and cycling of nitrogen in the Gulf of Maine.  
 959 *Journal of Marine Systems* **16**, 283–295.
- 960 TUCKER, M.A., BÖHNING-GAESE, K., FAGAN, W.F., FRYXELL, J.M., VAN MOORTER, B.,  
 961 ALBERTS, S.C., ALI, A.H., ALLEN, A.M., ATTIAS, N., AVGAR, T., BARTLAM-BROOKS,  
 962 H., BAYARBAATAR, B., BELANT, J.L., BERTASSONI, A., BEYER, D., ET AL. (2018)  
 963 Moving in the Anthropocene: global reductions in terrestrial mammalian  
 964 movements. *Science* **359**, 466–469.
- 965 VARPE, Ø., FIKSEN, Ø. & SLOTT, A. (2005) Meta-ecosystems and biological  
 966 energy transport from ocean to coast: the ecological importance of  
 967 herring migration. *Oecologia* **146**, 443–451.
- 968 VILLEGAS-RÍOS, D., RÉALE, D., FREITAS, C., MOLAND, E. & OLSEN, E.M. (2017)  
 969 Individual level consistency and correlations of fish spatial behaviour  
 970 assessed from aquatic animal telemetry. *Animal Behaviour* **124**, 83–94.

- 971 VITOUSEK, P.M. (2004) *Nutrient cycling and limitation: Hawaii as a model*  
972 *system*. Princeton University Press, Princeton.
- 973 VITOUSEK, P.M., MENGE, D.N.L., REED, S.C. & CLEVELAND, C.C. (2013) Biological  
974 nitrogen fixation: rates, patterns and ecological controls in terrestrial  
975 ecosystems. *Philosophical transactions of the Royal Society of London.*  
976 *Series B, Biological sciences* **368**, 20130119.
- 977 WAGNER, D., JONES, J.B. & GORDON, D.M. (2004) Development of harvester ant  
978 colonies alters soil chemistry. *Soil Biology and Biochemistry* **36**, 797–  
979 804.
- 980 WHILES, M.R., CALLAHAM, JR., M.A., MEYER, C.K., BROCK, B.L. & CHARLTON, R.E.  
981 (2001) Emergence of periodical cicadas (*Magicicada cassini*) from a  
982 Kansas riparian forest: densities, biomass and nitrogen flux. *American*  
983 *Midland Naturalist* **145**, 176–187.
- 984 WILLIAMS, J.J., PAPASTAMATIOU, Y.P., CASELLE, J.E., BRADLEY, D. & JACOBY, D.M.P.  
985 (2018) Mobile marine predators : an understudied source of nutrients to  
986 coral reefs in an unfished atoll. *Proceedings of the Royal Society of*  
987 *London B: Biological Sciences* **285**, 20172456.
- 988 WITHERINGTON, B.E. (1986) Human and natural causes of marine turtle clutch  
989 and hatchling mortality and their relationship to hatchling production on  
990 an important Florida nesting beach. Thesis. University of Central Florida,  
991 Orlando, Florida, USA.
- 992 WORM, B., BARBIER, E., BEAUMONT, N., DUFFY, J.E., FOLKE, C., HALPERN, B.S.,  
993 JACKSON, J.B.C., LOTZE, H., MICHELI, F., PALUMBI, S.R., SALA, E., SELKOE, K.,  
994 STACHOWICZ, J. & WATSON, R. (2006) Impacts of biodiversity loss on ocean  
995 ecosystem services. *Science* **314**, 787–790.
- 996 YANG, L.H. (2004) Periodical cicadas as resource pulses in North American  
997 forests. *Science* **306**, 1565–1567.
- 998 YANG, L.H. & GRATTON, C. (2014) Insects as drivers of ecosystem processes.  
999 *Current Opinion in Insect Science* **2**, 26–32.
- 1000 YOUNG, H.S., MCCAULEY, D.J., DUNBAR, R.B. & DIRZO, R. (2010) Plants cause  
1001 ecosystem nutrient depletion via the interruption of bird-derived spatial  
1002 subsidies. *Proceedings of the National Academy of Sciences* **107**, 2072–  
1003 2077.
- 1004 ZHAO, M. & RUNNING, S.W. (2010) Drought-induced reduction in global  
1005 terrestrial net primary production from 2000 through 2009. *Science* **329**,  
1006 940–943.

1007  
1008

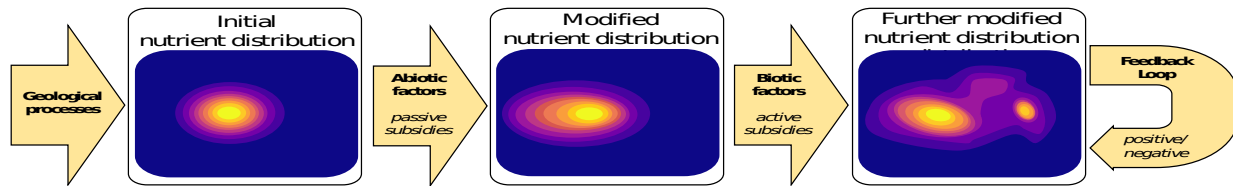


1009Table 1. Examples of behaviours that highlight various active subsidy  
1010features.

<b>Nature of active subsidy</b>	<b>Behaviour</b>	<b>Examples</b>	<b>References</b>
Counter-gradient transport	Migration/dispersal	Salmon spawning upstream	Helfield & Naiman (2001); Gende <i>et al.</i> (2002); Holtgrieve & Schindler (2011)
	Migration	Sea turtles migrating to nesting grounds, bringing ocean nutrients onto land	Bouchard & Bjørndal (2000)
	Foraging	'Whale pump' (see Fig. 4)	Roman & McCarthy (2010)
	Foraging	Pelagic birds foraging off islands, bringing ocean nutrients onto land	Anderson & Polis (1999); Adame <i>et al.</i> (2015); McFadden <i>et al.</i> (2016)
	Foraging	Hippopotamus daily foraging forays	Subalusky <i>et al.</i> (2015)
	Foraging	Reef sharks foraging in offshore waters	Williams <i>et al.</i> (2018)
Responding to resources	Habitat selection	Seabirds nesting on islands with suitable vegetation	Adame <i>et al.</i> (2015); Young <i>et al.</i> (2010); Graham <i>et al.</i> (2018)
	Foraging	Ungulates utilizing hotspots fertilized by their activity	Augustine (2003) McNaughton <i>et al.</i> (1997)
	Foraging	'Whale pump' (see Fig. 4)	Roman & McCarthy (2010)
	Habitat modification	Damselfish farming	Polunin & Koike, (1987); Hata <i>et al.</i> (2002)
Interacting with other agents	Habitat selection	Shoaling fish	McIntyre <i>et al.</i> (2008); Capps & Flecker (2013)
	Habitat selection	Central-place foraging in colonial ants	Clay <i>et al.</i> (2013)
	Anti-predator response	Changes in grasshopper foraging behaviours	Strickland <i>et al.</i> (2013)
	Anti-predator response	Alteration of ungulate space use	Frank (2008)
	Anti-predator response	Bats in caves during the day	Bird <i>et al.</i> (2007); Kunz <i>et al.</i> (2011)
	Parasite manipulation	Invertebrates jumping into streams	Sato <i>et al.</i> (2011)



1013

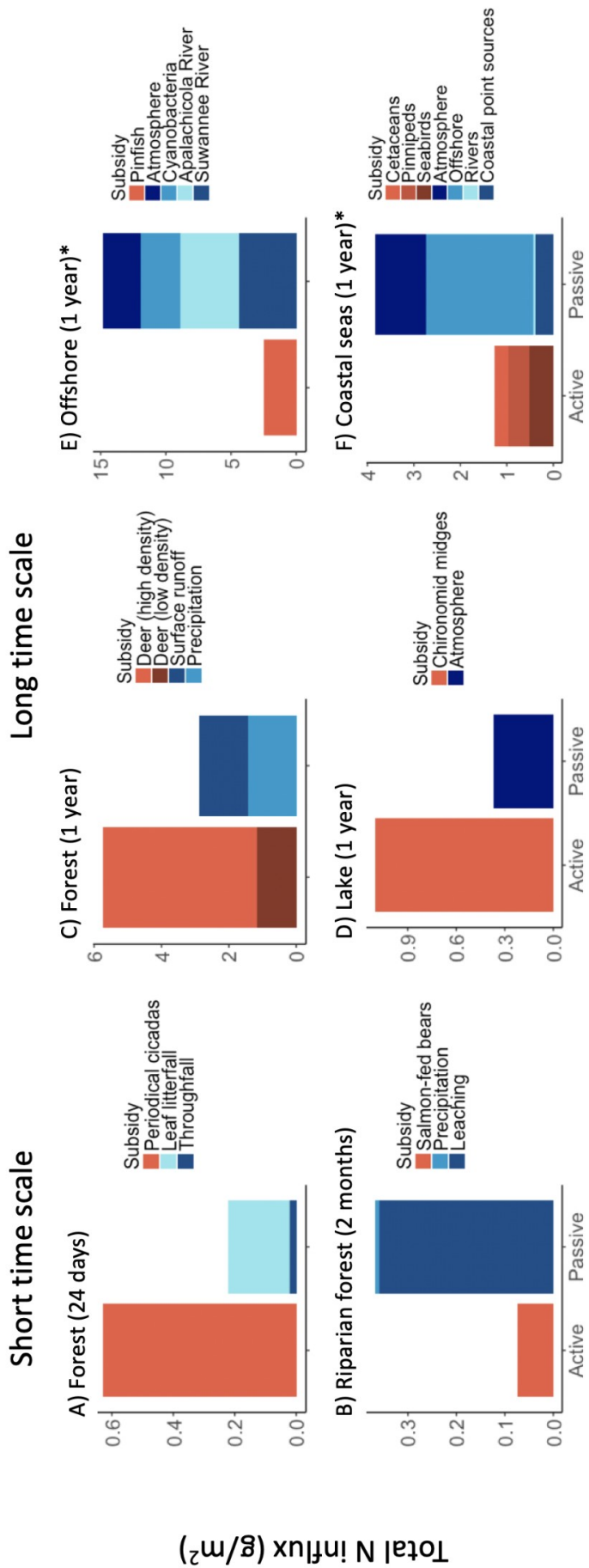


1014

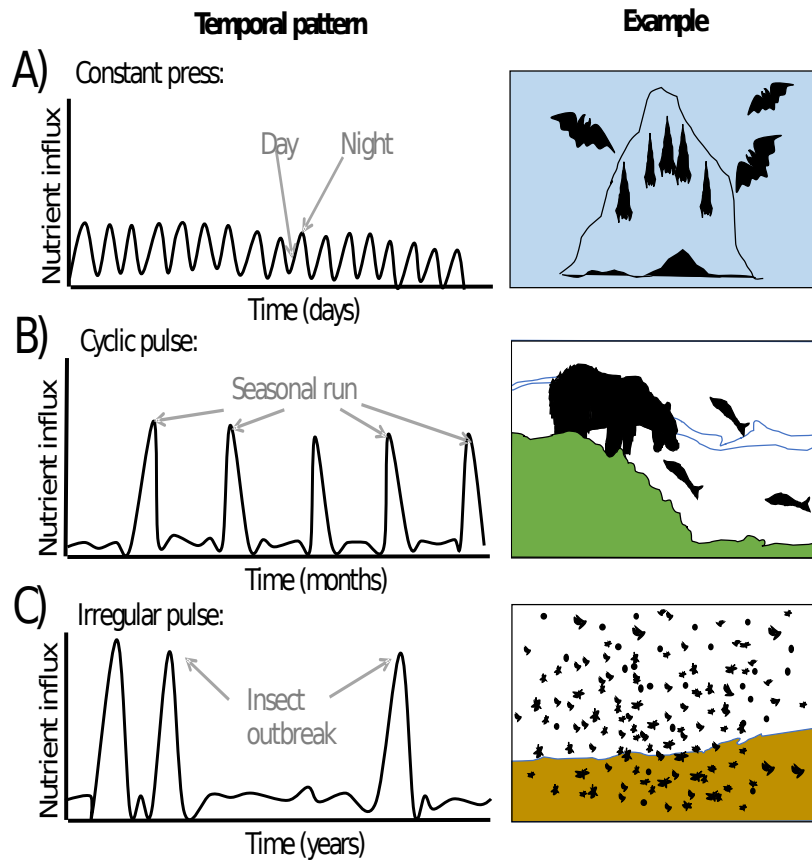
1015**Fig. 1.** The main forces shaping nutrient distributions. Initial distribution is a  
1016by-product of geological processes such as plate tectonics, volcanism and  
1017sedimentation (here, a hypothetical contour map, with warmer colours  
1018corresponding to higher concentrations of a certain nutrient, e.g. nitrogen or  
1019phosphorus). This distribution is reshaped by abiotic factors like erosion,  
1020ocean currents, wind, and gravity (here, creating an anisotropic peak shape).  
1021Animals respond to this distribution and further modify it by moving across  
1022local and regional scales. These movements often drive further  
1023heterogeneity (here, visualized as secondary peaks) at consistent hotspots of  
1024biological attractions or during pulses of nutrient flow such as at avian  
1025roosting colonies and in locust mass outbreaks. Finally, if animals respond  
1026continuously to resource distributions and the presence and behaviour of  
1027other organisms, this can generate feedback loops that further enhance  
1028heterogeneity.

1029

1030



**Fig. 2.** A quantitative comparison of net nitrogen influx by active (red) and passive (blue) subsidy sources in different ecosystems at various time scales. Active subsidies can generate pulses of events at shorter time scales: (A) in forest ecosystems, over the course of 24 days (Hamburg & Lin, 1998; Whiles *et al.*, 2001) and (B) in riparian forest ecosystems, over the course of 2 months (Helfield & Naiman, 2006). Similarly, they can create a prolonged impact over the course of an entire year: (C) in forest ecosystems (Peterjohn & Correll, 1984; Seagle, 2003); (D) along the aquatic-terrestrial interface in lake systems (Yang & Gratton, 2014; Dreyer *et al.*, 2015); (E) in offshore regions (Nelson *et al.*, 2013); and (F) in coastal marine ecosystems (Sowles, 2001; Townsend, 1998; Roman & McCarthy, 2010). In cases where a range of nitrogen-deposition values are available, we have included the most conservative estimates. Asterisks indicate log-transformed values.



1048

1049**Fig. 3.** Variation in temporal dynamics of nutrient influx by active subsidies.

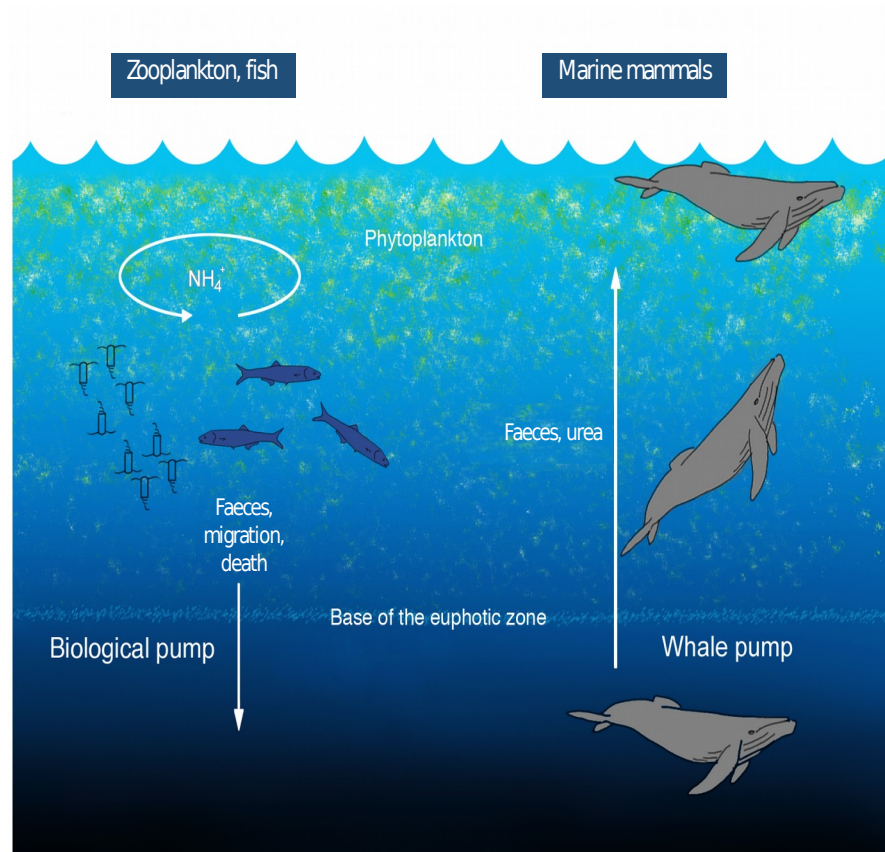
1050(A) Constant press: bats provide relatively constant subsidies (with a daily  
 1051cycle). These subsidies support a community of guano-dependent consumers  
 1052and their predators. (B) Cyclic pulse: migrating salmon provide strong pulses  
 1053of nutrients with an accurate yearly cycle. These pulses support upstream  
 1054consumers and facilitate aggregations of generally solitary bears. (C)  
 1055Irregular pulse: locust migrations provide massive but non-cyclic active  
 1056subsidies that support communities in arid areas.

1057

1058

1059

1060



1061

**Fig. 4.** The conceptual model of the whale pump (as presented by Roman & McCarthy, 2010). Great whales exhibit the key characteristics of active subsidy transport during their annual migrations, when they recycle nitrogen from deeper waters into the photic zone through a mechanism known as the 'whale pump' (Roman & McCarthy, 2010). Cetaceans feed at depth (>100 m) on fish and zooplankton in the waters through which they migrate, but must rise to the surface to breathe, releasing nitrogen-rich urea and faecal plume material. Nutrients from the waste (i.e. nitrogen) are utilized in the photic zone for growth and photosynthesis by phytoplankton, which are then consumed by zooplankton and fish to fuel the biological pump. This serves as



1072a counter-directional vector to the downward flux of key nutrients from the  
1073surface through the gravitational pull on aggregates and faeces, and the  
1074vertical movement of zooplankton and fish (Roman & McCarthy, 2010).



1075

1076**Fig. 5.** Behavioural features of agents that affect active subsidies. Marine  
1077birds act as a major transportation agent, bringing nutrients from the ocean  
1078into terrestrial systems on certain islands. The release of ocean-derived  
1079nutrients by birds is shown with orange arrows. This system demonstrates  
1080three general features that make active subsidies (animal-mediated nutrient  
1081transport) distinct from passive subsidies. (1) Birds transport nutrients  
1082counter-directionally to gravity, beyond the reach of ocean waves. (2) Birds  
1083respond to the nutrient distributions they generate, creating a positive-  
1084feedback loop. For example, on mangrove islands like that pictured here,  
1085nutrient influxes *via* guano at bird-aggregation sites create a nutrient

1086hotspot and facilitate growth of suitable roosting trees, which leads to  
1087positive feedback for bird populations (Adame *et al.*, 2015). (3) Transport is  
1088also influenced by interactions between conspecifics, creating a positive-  
1089feedback loop. Many seabirds use social information to inform their  
1090movement decisions, likely using the presence of conspecifics as a signal of  
1091high-quality nesting habitat (Forbes & Kaiser, 1994; see text for details).  
1092